

HIV CTL Epitopes

Following the entry for a given CTL epitope are brief comments explaining the context in which the epitope was defined. If the same epitope was studied in several labs, each study is cited in its own bulleted entry.

B. HIV CTL EPTOPES SORTED BY HLA RESTRICTING ELEMENT

This section presents tables of the epitopes included in Section A that have known HLA restricting elements, organized by the restricting element. Anchor and auxiliary residues for HLA molecules are listed, and if anchor residues with appropriate spacing are evident in the epitope, they are emboldened and underlined. This table provides minimal information about the epitopes, and only the shortest version of overlapping epitopes; for more information see the tables where epitopes are organized by protein location.

C. HIV PROTEIN EPTOPE MAPS

Because of the increasing number of defined epitopes, only human CTL and primate epitopes defined to within a region of 21 amino acids or less, with a known HLA specificity, are indicated on the HIV protein epitope maps.

The location and HLA restriction elements of CTL epitopes are indicated on protein sequences of the WEAU clone 1.60. These maps are meant to provide the relative location of epitopes on a given protein, but the WEAU sequence may not actually carry the epitope of interest, as it may vary relative to the sequence for which the epitope was defined. Epitopes are numbered in bold on the maps; the map numbering corresponding to the numbering of the epitope sequence alignments.

D. ALIGNMENTS

As with the maps, only human CTL epitopes defined within a region of 21 amino acids or less, with a known HLA specificity, have correspond-

ing alignments. For each numbered epitope in the epitope-protein maps, an alignment was generated from the protein sequence alignments in the HIV-1 genetic sequence database. All epitopes are aligned to the WEAU sequence and the sequence used to define the epitope is indicated directly above it. In consensus sequences an upper case letter indicates the amino acid was present in all sequences, a lower case letter indicates the amino acid was present in most sequences in a given position, and a question mark indicates two or more amino acids were represented with equal frequency. The master alignment files from which the epitope alignments were created are available at http://hiv-web.lanl.gov/ALIGN_98/ALIGN-INDEX-98.html. Included in the epitope alignments are only those sequences which completely span the gene of interest. Short fragments of sequences are excluded. The subtype designation and the country of isolation are indicated along with the common name of the sequence.

Subtype	Country of origin (Uganda)	Common name
A.	UG.	U455

The alignments were modified in some cases to optimize the alignment relative to the defined epitope and minimize insertions and deletions. A dash indicates identity to the consensus sequence, and a period indicates an insertion made to maintain the alignment. Stop codons are indicated with a \$, and frameshifts by a #; they are inserted to maintain the alignments. The alignments included in the printed version of this database contain only a subset of the sequences that are aligned in the immunology web site.

E. REFERENCES AND NOTES